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FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Aug 31, 2001 (20010831/UP).

=>

ANSWER 8 OF 10 USPATFULL L6

> For example, for treating or preventing chronic nonbacterial prostatitis, acute or chronic prostatitis, acute bacterial prostatitis, prostatodynia, congestive prostatitis, epididymitis, post-vasectomy pain and inflammation and/or urethritis in a patient a tachykinin receptor antagonist may be given in combination with such compounds as: an alpha blocker, especially an alpha-la blocker, such as doxazosin, indoramin, prazosin, tamsulosin, or terazosin; a 5-alpha reductase inhibitor, such as dutasteride or finasteride, especially a type 2 5-alpha reductase inhibitor, a dual 5-alpha reductase inhibitor, or combinations of type 1 and type 2 5-alpha reductase inhibitor; a prostate specific antigen conjugate; an antibiotic, including amikacin, amoxicillin, ampicillin, carbenicillin, cefaclor, cefadroxil, cefamandole, cefazolin, cefoxitin, cephalexin, cephalothin, cephapirin, cephradine, ciprofloxacin, cotrimoxazole, demeclocycline, doxycycline, erythromycin, gentamicin, kanamycin, methenamine hippurate, methenamine mandelate, minocycline, nalidixic acid, nitrofurantoin, norfloxacin, ofloxacin, sulfamethoxazole, sulfonamides, tetracycline, ticarcillin, tobramycin, trimethoprimin, or trimethoprimin-sulfamethoxazole, in particular a carbapenem antibiotic; anticholinergic agents, such as atropine, hyoscyamine, flavoxate, propantheline, or oxybutynin; a non-steroidal antiinflammatory, such as acetomeniphen, alprostadil, asprin, diclofenac, etodolac, ibuprofen, indomethacin, ketoprofe, ketorolac tromethamine, misoprostol, nabumetone, naproxen, naproxen sodium, oxaprozin, piroxicam, spironolactone, spironolactone with hydrochlorothiazide, or trovafloxacin; a corticosteroid; a selective cyclooxygenase-2 inhibitor, such as celecoxib, parecoxib, rofecoxib, valdecoxib, meloxicam, flosulide, nimesulide, MK-663, NS 398, DuP 697, SC-58125, SC-58635, or RS 57067; or a topical urinary analgesic, such as phenazopyridine, and salts thereof, and combinations thereof, and the like, as well as admixtures and combinations thereof.

ACCESSION NUMBER:

2000:50705 USPATFULL

TITLE:

SUMM

Method for treating or preventing chronic nonbacterial

prostatitis and prostatodynia

INVENTOR(S):

Guess, Harry A., Chapel Hill, NC, United States

Waldstreicher, Joanne, Scotch Plains, NJ, United States

Pearson, Jay Dee, Hatfield, PA, United States

PATENT ASSIGNEE(S): Merck & Co., Inc., Rahway, NJ, United States (U.S.

corporation)

NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.:

US 1999-313002 20000425 19990517 (9)

> NUMBER DATE

PRIORITY INFORMATION: US 1998-85866P 19980515 (60)

Utility

DOCUMENT TYPE: FILE SEGMENT:

Granted

PRIMARY EXAMINER:

Henley, III, Raymond

LEGAL REPRESENTATIVE:

Thies, J. Eric, Rose, David L.

NUMBER OF CLAIMS:

10

EXEMPLARY CLAIM:

CAPLUS COPYRIGHT 2003 ACS

50-55-5, Reserpine 50-60-2, Phentolamine 55-65-2, Guanethidine

55-73-2, Bethanidine 59-41-6, Bretylium 59-42-7, Phenylephrine

59-96-1, Phenoxybenzamine 59-98-3, Tolazoline 1131-64-2 4205-90-7,

Clonidine 19216-56-9, Prazosin

RL: BIOL (Biological study)

(sympathetically-maintained pain topical treatment

with)

ACCESSION NUMBER:

1991:623487 CAPLUS

DOCUMENT NUMBER:

115:223487

TITLE:

Compositions and methods of treatment of sympathetically-maintained pain using

.alpha.-adrenergic antagonists

INVENTOR(S):

Campbell, James N.

PATENT ASSIGNEE(S):

USA

SOURCE:

_ =>

PCT Int. Appl., 27 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT I	NO.		KI	DN	DATE				APPLICA	NOITA	NO.	DATE
MO	9112806			A1		19910905			WO 1991-US1318			318	19910226
	W: RW:	CA, AT.		CH.	DE.	DK.	ES.	FR.	GE	3, GR, 3	гт. т	U. NI	. SE
US	5070	•	,	A	,	1991	•	,		US 199	•	•	•
EP	EP 517850			Al 19921216			EP 1991-906357					19910226	
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GE	3, GR, :	IT, L	ıI, LU	, NL
JР	JP 05503539				2	1993	19930610			JP 199	1-506	069	19910226
JP	2786	538		B:	2	1998	0813						
US	5447	947		Α		1995	0905			US 1992	2-905	496	19920625
PRIORIT	Y APP	LN.	INFO.	:					US	1990-48	35156	5	19900226
									US	1991-66	31554	:	19910226
									WO	1991-U	31318	3	19910226

```
ANSWER 5 OF 52 CAPLUS COPYRIGHT 2001 ACS
L5
     1997:354939 CAPLUS
AN
DN
     127:61054
     Effects of transmural field stimulation in isolated smooth muscle of
ΤI
human
     rectum and internal anal sphincter
     Glavind, E. B.; Forman, A.; Madsen, G.; Tottrup, A.
AU
     Dep. Obstetrics Gynecology, Dep. Surgery L, Univ. Hospital Aarhus
CS
Surgical
     Res. Unit,, Aarhus, DK-8000, Den.
     Am. J. Physiol. (1997), 272(5, Pt. 1), G1075-G1082
     CODEN: AJPHAP; ISSN: 0002-9513
PB
     American Physiological Society
DT
     Journal
     English
LA
CC
     2-8 (Mammalian Hormones)
     Smooth muscle prepns. from the circular muscle layer of the most distal
AB
     rectum and the proximal and distal human internal anal sphincter (IAS)
    mounted in organ baths to record isometric tension developed spontaneous
     tension. Transmural elec. field stimulation (TMS) induced frequency- and
    impulse duration-dependent relaxations sensitive to tetradotoxin in the
     stimulation range of 0.5-40 Hz and 0.04-0.6 ms. Poststimulus
contractions
     were most frequent and prominent in rectal prepns. Maximal relaxations
     were comparable in the three locations and were achieved at 10 Hz and 0.4
          The frequency inducing half-maximal response was lower in rectal
     strips compared with IAS. Phentolamine (10-6 M) enhanced relaxations and
     diminished off-contractions at 40 Hz in distal IAS. N.omega.-nitro-L-
     arginine (L-NNA) concn. dependently inhibited both relaxations and
     off-contractions (10 Hz, 0.4 ms). The pD2 values (-log E50) of L-NNA
were
     lower in rectal muscle compared with those in IAS. L-Arginine (10-4 M)
     inhibited the blocking effect of L-NNA. In one-half of the prepns.,
     reversed the relaxations to duration contractions (15-40 Hz), which were
     inhibited by atropine in rectal prepns. and by phentolamine in IAS. In
     conclusion, excitatory innervation of the IAS is .alpha.-adrenergic and
     cholinergic in the rectum. A product of the L-arginine-nitric oxide
     pathway mediates the TMS-induced inhibition of the muscle and is also
     involved in poststimulus contractions.
     nerve adrenergic cholinergic rectum analysis sphincter
ST
     Cholinergic neurons
IT
     Contraction (muscle)
     Neuromuscular transmission
     Rectum
     Smooth muscle
        (adrenergic and cholinergic regulation of transmural field stimulation
        effects in isolated smooth muscle of human rectum and internal anal
        sphincter)
     Intestine
IT
        (internal anal sphincter; adrenergic and cholinergic regulation of
        transmural field stimulation effects in isolated smooth muscle of
human
        rectum and internal anal sphincter)
IT
     Nervous system
        (.alpha.-adrenergic; adrenergic and
        cholinergic regulation of transmural field stimulation effects in
        isolated smooth muscle of human rectum and internal anal
```

sphincter)
IT 10102-43-9, Nitric oxide, biological studies
 RL: BAC (Biological activity or effector, except adverse); BPR
(Biological
 process); BIOL (Biological study); PROC (Process)
 (adrenergic and cholinergic regulation of transmural field stimulation effects in isolated smooth muscle of human rectum and internal anal sphincter)

```
L19 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2001 ACS
      Fecal incontinence and anal itch can be treated by administration, more
AΒ
      particularly by local application to the anus, of an .alpha.
      adrenergic blocker, nitric oxide synthase
      inhibitor, prostaglandin F2.alpha., dopamine, morphine, .beta.-
      blockers, and 5-Hydroxytryptamine. The patients who benefit most
      from the invention are those who have a normal or low max. anal resting
      pressure and a structurally intact internal anal
      sphincter muscle, and patients who have had major bowel resection
      and reanastomosis. Phenylephrine-Hcl was added to a base cream to form.
                             1998:479406 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                             129:86054
                             Pharmaceutical composition for treating fecal
TITLE:
                             Kamm, Michael Albert; Phillips Wobin Kenneth Stewart UK
PCT Int. Appl., 28 pp.
CODEN: PIXXP2
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
                             Patent
LANGUAGE:
                             English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                   APPLICATION NO. DATE
      PATENT NO.
                       KIND DATE
                         ____
                                 19980 702
                                                  WO 1997-GB3525
                                                                       19971223
      WO 9827971
                          A1
          W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
                                 19980717
                                                                       19971223
      AU 9853315
                          A1
                                                  AU 1998-53315
                                 20010118
      AU 728889
                           B2
                           A1
                                 19991006
                                                  EP 1997-950311
                                                                       19971223
      EP 946155
          R: AT, BE, CH, DE, DK ES, FR, GB, GR, IT, LI, NL, SE, PT, IE, FI
      JP 2001507020
                                 20010529
                                                   JP 1998-528550
                                                                       19971223
                          Т2
                                                                  A 19961223
PRIORITY APPLN. INFO .:
                                               GB 1996-26739
                                                                   A 19961223
                                               GB 1996-26750
                                                                   Α
                                               GB 1997-3309
                                                                       19970218
                                                                 W 19971223
                                               WO 1997-GB3525
     ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2001 ACS
L19
TI
      Nervous control of the internal anal sphincter of the
      cat
      Hypogastric nerve stimulation elicited slow time course depolarization
AΒ
      responses in anal sphincteric circular muscle of cats,
      which were abolished by .alpha.-adrenergic
      blockers. Stimulation of parasympathetic outflow to the internal
      anal sphincter (2nd ventral sacral root, VS2) inhibited
      spontaneous elec. activity of the circular muscle, apparently through
      intramural nonadrenergic, noncholinergic (purinergic) inhibitory
      neurons. Rectal distension also inhibited anal
      sphincteric circular muscle via nonadrenergic, noncholinergic
      intramural neurons. Longitudinal muscle responses to VS2 or hypogastric
      nerve stimulation indicated that the muscle receives excitatory
      innervation from preganglionic parasympathetic nerves connected with
      intramural cholinergic neurons, and inhibitory sympathetic
      innervation from noradrenergic axons running in the hypogastric nerves.
```

```
Responses of circular muscle to simultaneous VS2 and hypogastric nerve. .
Nervous system
```

by) ΙT Nervous system

ΙT

(sympathetic, anal sphincter muscle regulation by)

1981:100656 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 94:100656

Nervous control of the internal anal TITLE:

sphincter of the cat

(parasympathetic, anal sphincter muscle regulation

Bouvier, M.; Gonella, J. AUTHOR(S):

Dep. Neurophysiol. Veg., Inst. Neurophysiol. CORPORATE SOURCE:

> Psychophysiol., Marseille, 13274/2, Fr. J. Physiol. (London) (1981), 310 457-69

CODEN: JPHYA7; ISSN: 0022-3751

DOCUMENT TYPE:

SOURCE:

Journal English LANGUAGE:

L19 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2001 ACS

Effects of rectal distension on the internal anal sphincter of cats

The effect of i.v. autonomimetic drugs and blocking drugs on the AB contractions and relaxations of the internal anal sphincter were studied in anesthetized cats with a miniature air-filled intraluminal balloon placed in the middle 3rd of the rectum. Acetylcholine. . . by atropine. Noradrenaline [51-41-2] and adrenaline [51-43-4] produced biphasic responses of contraction and relaxation which were abolished by the .alpha.-receptor blocker dihydroergotamine tartrate. The relaxation response to isoprenaline [7683-59-2] was abolished by the .beta.-adrenergic blocker propranolol. prolonged relaxation of the internal sphincter occurred upon rectal distension which was abolished by the ganglion blocker hexamethonium. These expts. suggest that the tone of the internal anal sphincter is under complex neural control involving cholinergic and .alpha.-adrenergic motor pathways and .beta.-adrenergic and noncholinergic nonadrenergic inhibitory pathways. Reflex responses to rectal distension are influenced by all of these mechanisms.

ΙT 51-41-2 51-43-4 51-84-3 7683-59-2

RL: BIOL (Biological study)

(internal anal sphincter contraction in response

to)

ACCESSION NUMBER: 1972:522160 HCAPLUS

DOCUMENT NUMBER: 77:122160

TITLE: Effects of rectal distension on the internal

anal sphincter of cats

Garrett, J. R.; Howard, E. R. AUTHOR(S):

King's Coll. Hosp. Med. Sch., London, Engl.
J. Physiol. (London) (1972), 222(1), 85P-86P CORPORATE SOURCE: SOURCE:

CODEN: JPHYA7

DOCUMENT TYPE: Journal English LANGUAGE:

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SINCE FILE COST IN U.S. DOLLARS TOTAL ENTRY SESSION FULL ESTIMATED COST 30.46 143.21

SINCE FILE DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -1.76-2.32

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L19 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2001 ACS
AN
     1972:522160 HCAPLUS
DN
     77:122160
     Effects of rectal distension on the internal anal
ΤI
     sphincter of cats
ΑU
     Garrett, J. R.; Howard, E. R.
CS
     King's Coll. Hosp. Med. Sch., London, Engl.
     J. Physiol. (London) (1972), 222(1), 85P-86P
SO
     CODEN: JPHYA7
DΤ
     Journal
     English
LA
CC
     1-5 (Pharmacodynamics)
     The effect of i.v. autonomimetic drugs and blocking drugs on the
AΒ
     contractions and relaxations of the internal anal
     sphincter were studied in anesthetized cats with a miniature
     air-filled intraluminal balloon placed in the middle 3rd of the rectum.
     Acetylcholine [51-84-3] caused sphincter contraction which was blocked by
     atropine. Noradrenaline [51-41-2] and adrenaline [51-43-4] produced
     biphasic responses of contraction and relaxation which were abolished by
     the .alpha.-receptor blocker dihydroergotamine tartrate. The
     relaxation response to isoprenaline [7683-59-2] was abolished by the
     .beta.-adrenergic blocker propranolol. A prolonged relaxation
     of the internal sphincter occurred upon rectal distension which was
     abolished by the ganglion blocker hexamethonium. These expts.
     suggest that the tone of the internal anal sphincter
     is under complex neural control involving cholinergic and .alpha
     .-adrenergic motor pathways and .beta.-adrenergic and
     noncholinergic nonadrenergic inhibitory pathways. Reflex
     responses to rectal distension are influenced by all of these mechanisms.
ST
     sphincter contraction adrenergic; cholinergic sphinter contraction;
     adrenaline rectal distension; noradrenaline rectal distension;
     acetylcholine rectal distension
ΙT
     Intestine
        (sphincter anae, autonomic control of)
ΙT
     51-41-2
             51-43-4
                         51-84-3
                                   7683-59-2
     RL: BIOL (Biological study)
        (internal anal sphincter contraction in response
        to)
```

```
L19 ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2001 ACS
     1981:100656 HCAPLUS
AN
DN
     94:100656
     Nervous control of the internal anal sphincter of the
ΤI
ΑU
     Bouvier, M.; Gonella, J.
     Dep. Neurophysiol. Veg., Inst. Neurophysiol. Psychophysiol., Marseille,
CS
     13274/2, Fr.
     J. Physiol. (London) (1981), 310 457-69
SO
     CODEN: JPHYA7; ISSN: 0022-3751
DT
     Journal
     English
LA
CC
     13-13 (Mammalian Biochemistry)
     Hypogastric nerve stimulation elicited slow time course depolarization
AΒ
     responses in anal sphincteric circular muscle of cats,
     which were abolished by .alpha.-adrenergic
     blockers. Stimulation of parasympathetic outflow to the internal
     anal sphincter (2nd ventral sacral root, VS2) inhibited
     spontaneous elec. activity of the circular muscle, apparently through
     intramural nonadrenergic, noncholinergic (purinergic) inhibitory
     neurons. Rectal distension also inhibited anal
     sphincteric circular muscle via nonadrenergic, noncholinergic
     intramural neurons. Longitudinal muscle responses to VS2 or hypogastric
     nerve stimulation indicated that the muscle receives excitatory
     innervation from preganglionic parasympathetic nerves connected with
     intramural cholinergic neurons, and inhibitory sympathetic
     innervation from noradrenergic axons running in the hypogastric nerves.
     Responses of circular muscle to simultaneous VS2 and hypogastric nerve
     stimulation indicated that the release of noradrenaline from sympathetic
     nerves is modulated by muscarinic and nicotinic receptors located on
     noradrenergic nerve endings, which abolish and increase release, resp.
     anus sphincter muscle innervation; nerve parasympathetic sympathetic anus
ST
     sphincter; receptor anus sphincter muscle
ΙT
     Receptors
     RL: PROC (Process)
        (of anus sphincter muscle, characterization of)
ΙT
        (anus, sphincter, parasympathetic and sympathetic nervous control of)
ΙT
     Nervous system
        (parasympathetic, anal sphincter muscle regulation
ΙT
     Nervous system
        (sympathetic, anal sphincter muscle regulation by)
```

L19 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2001 ACS

TI Effects of rectal distension on the internal anal sphincter of cats

The effect of i.v. autonomimetic drugs and blocking drugs on the ΑŔ contractions and relaxations of the internal anal sphincter were studied in anesthetized cats with a miniature air-filled intraluminal balloon placed in the middle 3rd of the rectum. Acetylcholine. . . by atropine. Noradrenaline [51-41-2] and adrenaline [51-43-4] produced biphasic responses of contraction and relaxation which were abolished by the .alpha.-receptor blocker dihydroergotamine tartrate. The relaxation response to isoprenaline [7683-59-2] was abolished by the .beta.-adrenergic blocker propranolol. A prolonged relaxation of the internal sphincter occurred upon rectal distension which was abolished by the ganglion blocker hexamethonium. These expts. suggest that the tone of the internal anal sphincter is under complex neural control involving cholinergic and .alpha.-adrenergic motor pathways and .beta.-adrenergic and noncholinergic nonadrenergic inhibitory pathways. Reflex responses to rectal distension are influenced by all of these mechanisms.

Ļ

IT 51-41-2 51-43-4 51-84-3 7683-59-2

RL: BIOL (Biological study)

(internal anal sphincter contraction in response

to)

ACCESSION NUMBER: 1972:522160 HCAPLUS

DOCUMENT NUMBER: 77:122160

TITLE: Effects of rectal distension on the internal

anal sphincter of cats

AUTHOR(S): Garrett, J. R.; Howard, E. R.

CORPORATE SOURCE: King's Coll. Hosp. Med. Sch., London, Engl. SOURCE: J. Physiol. (London) (1972), 222(1), 85P-86P

CODEN: JPHYA7

DOCUMENT TYPE: Journal LANGUAGE: English

L34 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2001 ACS 1981:100656 CAPLUS AN 94:100656 DN Nervous control of the internal anal sphincter of the cat ΤI Bouvier, M.; Gonella, J. ΑU Dep. Neurophysiol. Veg., Inst. Neurophysiol. Psychophysiol., Marseille, CS 13274/2, Fr. SO J. Physiol. (London) (1981), 310 457-69 CODEN: JPHYA7; ISSN: 0022-3751 DT Journal LA English 13-13 (Mammalian Biochemistry) CC Hypogastric nerve stimulation elicited slow time course depolarization AB responses in anal sphincteric circular muscle of cats, which were abolished by .alpha.-adrenergic blockers. Stimulation of parasympathetic outflow to the internal anal sphincter (2nd ventral sacral root, VS2) inhibited spontaneous elec. activity of the circular muscle, apparently through intramural nonadrenergic, noncholinergic (purinergic) inhibitory neurons. Rectal distension also inhibited anal sphincteric circular muscle via nonadrenergic, noncholinergic intramural neurons. Longitudinal muscle responses to VS2 or hypogastric nerve stimulation indicated that the muscle receives excitatory innervation from preganglionic parasympathetic nerves connected with intramural cholinergic neurons, and inhibitory sympathetic innervation from noradrenergic axons running in the hypogastric nerves. Responses of circular muscle to simultaneous VS2 and hypogastric nerve stimulation indicated that the release of noradrenaline from sympathetic nerves is modulated by muscarinic and nicotinic receptors located on noradrenergic nerve endings, which abolish and increase release, resp. anus sphincter muscle innervation; nerve parasympathetic sympathetic anus ST sphincter; receptor anus sphincter muscle ΙT Receptors RL: PROC (Process) (of anus sphincter muscle, characterization of) Intestine IT (anus, sphincter, parasympathetic and sympathetic nervous control of) Nervous system TΤ (parasympathetic, anal sphincter muscle regulation by) ΙT Nervous system

(sympathetic, anal sphincter muscle regulation by)

L24 ANSWER 19 OF 20 HCAPLUS COPYRIGHT 2001 ACS

AN 1997:74661 HCAPLUS

DN 126:152610

TI Effects of loperamide oxide on gastrointestinal transit time and anorectal function in patients with chronic diarrhea and **fecal**incontinence

AU Sun, W. M.; Read, N. W.; Verlinden, M.

CS Royal Adelaide Hospital, Adelaide, Australia

SO Scand. J. Gastroenterol. (1997), 32(1), 34-38

CODEN: SJGRA4; ISSN: 0036-5521

PB Scandinavian University Press

DT Journal

LA English

CC 1-9 (Pharmacology)

Loperamide improves anorectal function in patients with chronic diarrhea. AB We wished to investigate whether the prodrug loperamide oxide has similar effects. Eleven patients with chronic diarrhea and fecal incontinence participated in a randomized, placebo-controlled, double-blind, crossover study of the effects of loperamide oxide (4 mg twice daily for 1 wk). Loperamide oxide reduced wet stool wt. and improved the patients' ratings of symptoms. Mouth-to-cecum transit time was not altered, but whole-gut transit time was prolonged. There were limited effects on anorectal function, but the mean min. basal pressure mainly contributed by the internal anal sphincter (IAS) was increased, as was the mean vol. infused before leakage occurred in the saline continence test. Loperamide oxide is effective in the treatment of diarrhea with fecal incontinence; normalization of colon transit time and an increase in the tone of the IAS seem to be the main determinants of efficacy.

ST loperamide oxide antidiarrheal

IT Antidiarrheals

(effects of loperamide oxide on gastrointestinal transit time and anorectal function in humans with chronic diarrhea and **fecal** incontinence)

IT 106900-12-3, Loperamide oxide

incontinence)

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (effects of loperamide oxide on gastrointestinal transit time and anorectal function in humans with chronic diarrhea and **fecal**

A CONTRACTOR OF THE PARTY OF TH

L34 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2001 ACS

AN 1999:186986 CAPLUS

DN 131:16909

TI Membrane properties and the neuro-effector transmission of smooth muscle cells in the canine internal **anal** sphincter

AU Kubota, Masayuki; Suita, Sachiyo; Szurszewski, Joseph H.

CS Department of Pediatric Surgery Faculty of Medicine, Kyushu University, Fukuoka, 812-8582, Japan

SO J. Smooth Muscle Res. (1998), 34(4), 173-184 CODEN: JSMRE2; ISSN: 0916-8737

PB Japanese Society of Smooth Muscle Research

DT Journal

LA English

CC 13-6 (Mammalian Biochemistry)

AB The most distal part of the circular muscle layer functions as the internal anal sphincter, which constitutes a high

pressure zone at rest, but maintains a relaxed state during defecation. To elucidate such sphincter mechanisms of the smooth muscle cells, the circular muscle layer in the canine anal canal was examd. within 2 cm from the anal verge. Both the mech. and intracellular elec. activities were recorded simultaneously. The examd. region could be divided into three different regions according to the pattern of spontaneous activity and innervation and consisted of an upper region (20-15 mm from the anal verge), a transitional region (15-5 mm from the anal verge) and a lower region (within 5 mm from the anal verge), resp. The spontaneous membrane activity was characterized by ongoing slow potential changes and each potential change was assocd. with a phasic contraction in the three regions. The mean frequencies of spontaneous elec. activity were 6.8, 15.9, and 24.1 c/min in the upper, transitional and lower regions, resp. In the transitional and lower region, muscle tone generation was obsd. Transmural field stimulation (0.4 ms in pulse duration) evoked membrane depolarization and contractions in the lower region. The application of an .alpha.-adrenergic blocking agent,

completely suppressed the generation of excitatory responses, leaving a long-lasting hyperpolarization assocd. with relaxation. In the transitional and upper region, stimulation consistently evoked membrane hyperpolarization with relaxation. The characteristics of this hyperpolarization response varied among the three regions. The total duration of hyperpolarization increased distally, while the time to peak-hyperpolarization became decreases in a reverse direction. These regional differences in the characteristics of spontaneous membrane activity and innervation indicate that the transitional and lower region might therefore function as the internal anal sphincter

ST smooth muscle internal sphincter membrane property neurotransmission IT Membrane potential

(biol.; membrane properties and the neuro-effector transmission of smooth muscle cells in the canine internal **anal** sphincter)

IT Polarization

(hyperpolarization, biol.; membrane properties and the neuro-effector transmission of smooth muscle cells in the canine internal **anal** sphincter)

IT Intestine

(internal anal sphincter; membrane properties and the neuro-effector transmission of smooth muscle cells in the canine internal anal sphincter)

IT Cell membrane

Muscle contraction

(membrane properties and the neuro-effector transmission of smooth muscle cells in the canine internal **anal** sphincter)

IT Muscle

(smooth; membrane properties and the neuro-effector transmission of smooth muscle cells in the canine internal anal sphincter)

RE.CNT 21

RE

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- (2) Boeckxstaens, G; Br J Pharmacol 1993, V109, P1079 CAPLUS
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ANSWER 15 OF 20 HCAPLUS COPYRIGHT 2001 ACS
ΑN
     1998:479406 HCAPLUS
DN
     129:86054
ΤI
     Pharmaceutical composition for treating fecal
     incontinence and anal itch
IN
     Kamm, Michael Albert; Phillips, Robin Kenneth Stewart
PA
     PCT Int. Appl., 28 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     English
LA
     ICM A61K031-00
IC
     ICS A61K031-135; A61K031-485; A61K031-195; A61K031-557; A61K031-40
CC
     63-6 (Pharmaceuticals)
FAN.CNT 1
                                           ` APPLICATION NO. DATE
                      KIND DATE
     PATENT NO.
     ______
                                              _____
     WO 9827971 A1 19980702
                                            WO 1997-GB3525 19971223
ΡI
         W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
              DK, EE, ES, FI, GB, GE, GH, GW, HU, ID, IL, IS, JP, KE, KG, KP,
              KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO,
              NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA,
              UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI,
              FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM,
              GA, GN, ML, MR, NE, SN, TD, TG
                                              AU 1998-53315
                                                                19971223
     AU 9853315
                       A1
                              19980717
     AU 728889
                        В2
                              20010118
                                             EP 1997-950311 19971223
     EP 946155
                        A1
                              19991006
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, SE, PT, IE, FI
                    T2 20010529
A 19961223
A 19961223
A 19970218
                                            JP 1998-528550 19971223
     JP 2001507020
PRAI GB 1996-26739
     GB 1996-26750
     GB 1997-3309
     WO 1997-GB3525 W
                              19971223
     Fecal incontinence and anal itch
AB
     can be treated by administration, more particularly by <del>local application</del>
     to the anus, of an alpha. adrenergic blocker, mitric oxide synthase inhibitor, prostaglandin F2.alpha., dopamine, morphine, beta.-blockers, and 5-Hydroxytryptamine. The patients who benefit most from the invention are those who have a normal or low max. anal resting pressure and a
     structurally intact internal anal sphincter muscle, and patients
     who have had major bowel resection and reanastomosis. Phenylephrine-HCl
     was added to a base cream to form a compn.
     pharmaceutical fecal incontinence anus itch
ST
ΙT
     Intestine
         (anus; pharmaceutical compn. for treating fecal
        incontinence and anal itch)
ΙT
     Drug delivery systems
         (foams; pharmaceutical compn. for treating fecal
        incontinence and anal itch)
     Feces
ΤT
     Ointments (drug delivery systems)
     Sprays (drug delivery systems)
     Suppositories (drug delivery systems)
     Suspensions (drug delivery systems)
     .alpha.1-Adrenoceptor agonists _
     .beta.-Adrenoceptor antagonists
         (pharmaceutical compn. for treating fecal
        incontinence and anal itch)
ΙT
     125978-95-2, Nitric oxide synthase
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
```

:: 1.

(inhibitors; pharmaceutical compn. for treating **fecal** incontinence and anal itch)

IT 50-67-9, 5-Hydroxytryptamine, biological studies 51-41-2, Noradrenaline 51-61-6, Dopamine, biological studies 57-27-2, Morphine, biological studies 59-42-7, Phenylephrine 61-76-7, Phenylephrine hydrochloride 390-28-3, Methoxamine 551-11-1, Prostaglandin F2.alpha. 2149-70-4, L-Ornithine, N5-[imino(nitroamino)methyl]- 35700-23-3, Carboprost 50903-99-6, L-NAME

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (pharmaceutical compn. for treating fecal
 incontinence and anal itch)

: - 4.

L28 ANSWER 5 OF 510 CAPLUS COPYRIGHT 2001 ACS

. . . system in acclimatizing to high altitude in men. The purpose of this investigation was to det. the extent to which .alpha.adrenergic blockade affects the sympathoadrenal responses to exercise during acute high-altitude exposure in women. Twelve eumenorrheic women (24.7.+-.1.3 yr, 70.6.+-.2.6 kg). . at sea level (on sep. days) on a bicycle ergometer after 3 days of taking either a placebo or an .alpha.-blocker (3 mg/day prazosin). Subjects also performed two similar exercise tests while at altitude. Effectiveness of blockade was detd. by phenylephrine challenge. At sea level, plasma norepinephrine levels during exercise were 48% greater when subjects were .alpha.-blocked compared with their placebo. . . obsd. for plasma epinephrine levels during exercise. No phase differences were obsd. across any condition studied. It was concluded that .alpha.-adrenergic blockade resulted in a compensatory sympathoadrenal response during exercise at sea level and altitude, and this effect was more pronounced. . .

'ANSWER 2 OF 2 CAPLUS COPYRIGHT 2001 ACS 1990:509846 CAPLUS

ΑN

DN113:109846

Role of alpha adrenoceptors in opossum internal anal sphincter ΤI

ΑU Yamato, Shigeru; Rattan, Satish

Div. Gastroenterol., Beth Israel Hosp., Boston, MA, 02215, USA CS

J. Clin. Invest. (1990), 86(2), 424-9

CODEN: JCINAO; ISSN: 0021-9738

Journal DT LA

SO

English

CC 2-8 (Mammalian Hormones)

AB The role of .alpha.-adrenoceptors in the internal anal sphincter (IAS) of opossum was studied. Resting pressure in the IAS (IASP) was recorded using low compliant continuously perfused catheters. The effects of the .alpha.1-adrenoceptor agonist phenylephrine and .alpha.2-adrenoceptor agonist clonidine and their corresponding selective antagonists, prazosin and yohimbine, resp., were examd. on the resting IASP, and on rectal balloon distension (RBD) -mediated IAS relaxation. Phenylephrine caused a rise in the IASP that was blocked by prazosin and not by yohimbine. Phenylephrine had no effect on IAS relaxation caused by RBD. Clonidine on the other hand caused significant suppression of IAS relaxation in response to RBD, but caused minimal changes in the resting IASP. The suppression of IAS relaxation by clonidine was selectively antagonized by yohimbine but not by prazosin. Thus, .alpha.2-adrenoceptors exert important neuromodulatory influences on rectoanal inhibitory reflex, whereas .alpha.1-adrenoceptors may exert modulatory effects on the resting IAS tone.

anus internal sphincter adrenergic receptor ST

Intestine-IT

(anus, internal sphincter, function of, adrenergic receptors regulation of)

ΙT Receptors

RL: BIOL (Biological study)

(.alpha.1-adrenergic, internal anal. sphincter function_regulation_by)_

IT Receptors

RL: BIOL (Biological study)

(.alpha.2-adrenergic, internal anal. sphincter function regulation by)

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